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(54) Title: TETRACYCLINE REPRESSOR-MEDIATED BINARY REGULATION SYSTEM FOR CONTROL OF GENE EXPRESSION IN TRANSGENIC ANIMALS

(57) Abstract

The present invention relates to a tetracycline repressor-mediated binary regulation system for the control of gene expression in transgenic animals. It is based, at least in part, on the discovery that, in a non-human transgenic animal that carries a first transgene under the control of a modified promoter comprising a tetR operator sequence and a second transgene encoding the tetR repressor protein, expression of the first transgene may be efficiently induced by administering tetracycline to the animal.

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TETRACYCLINE REPRESSOR-MEDIATED BINARY REGULATION SYSTEM FOR CONTROL OF GENE EXPRESSION IN TRANSGENIC ANIMALS

1. INTRODUCTION

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The present invention relates to a tetracycline repressor-mediated binary regulation system for the control of gene expression in transgenic animals. It is based, at least in part, on the discovery that, in a non-human transgenic animal that carries a first transgene under the control of a modified promoter comprising a tetR operator sequence and a second transgene encoding the tetR repressor protein, expression of the first transgene may be efficiently induced by administering tetracycline to the animal.

2. BACKGROUND OF THE INVENTION

2.1. CONTROL OF GENE EXPRESSION IN TRANSGENIC ANIMALS

20 The production of transgenic animals for both experiment and agricultural purposes is now well known (Wilmut et al., 7 July 1988, New Scientist pp. 56-59). In research, transgenic animals are a powerful tool that have made significant contributions to our 25 understanding of many aspects of biology and have contributed to the development of animal models for human diseases (Jaenisch, 1988, Science 240:1468-It is also clear that several livestock species can be made transgenic and these species 30 promise to expand and revolutionize the method of production and diversity of pharmaceutical products available in the future, in addition to improving the agricultural qualities of the livestock species (Wilmut et al., supra).

A critical, often neglected, aspect of developing transgenic animals is the process whereby expression

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of the newly introduced gene, referred to as the transgene, is controlled. This is an important process since stringent regulation of transgene expression is often important both for practical, regulatory and safety reasons and to maintain the health of the transgenic animal. In the past either "inducible" or "tissue specific" regulatory mechanisms have been used. Inducible regulation is defined herein as a method of gene regulation which allows for some form of outside manipulation of the onset and/or level of transgene expression. Tissue specific regulation is defined herein as a method for targeting transgene expression to particular tissues or organs.

Inducible gene regulation may be achieved using relatively simple promoter systems such as the metallothionein heat shock promoters, or by using promoters which are responsive to specific compounds such as the Mouse mammary tumor virus LTR which is responsive to glucocorticoid stimulation. 20 flexible, though more complex inducible regulation systems can be achieved through a "binary" gene approach which utilizes a transactivator gene product to control expression of a second gene of interest. Tissue specific gene regulation usually consists of 25 simple single gene methods (Byrne et al., 1989, Proc. Natl. Acad. Sci. U.S.A. 86:5473-5477; Ornitz et al., 1991, Proc. Natl. Acad. Sci. U.S.A. <u>88</u>:698-702), although binary transactivator systems can also provide a high degree of tissue specificity.

These current systems provide only a limited ability to control the time of transgene expression within individual animals. In this respect tissue specific promoter elements provide no method to control the onset of transgene activity, but function merely to target gene expression to defin d sites.

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Simple inducible promoters such as metallothionein generally lack tissue specificity and usually have some aspect of endogenous basal expression which cannot be controlled. Thus even for the extensively used inducible metallothionein promoter this approach at best only permits selection of the time at which a relative increase in transgene expression can be induced.

induced. Binary transactivation systems typically consist 10 of two transgenic animals. One animal contains the gene of interest controlled by a promoter element that requires a specific transactivator gene product for expression. Thus, the gene of interest is not expressed in the absence of the transactivator. 15 second transgenic animal is then made which expresses the required transactivator in the desired tissue. mating these two transgenic animals, offspring containing both the gene of interest and the transactivator transgene can be produced. 20 these doubly transgenic animals is the gene of interest expressed. Since expression of the gene of interest requires the transactivator, this binary approach dramatically reduces or eliminates any undesirable basal expression inherent in simple 25 inducible systems. Additionally, if expression of the transactivator is targeted using a tissue specific promoter, then in the double transgenics, expression of the gene of interest is in effect targeted to the same specific tissue. Binary systems provide 30 therefore a low resolution method of temporal regulation in as much as they allow the determination of which generation of animals will express the gene

35 however, to control the time and level of gene expression within an individual transgenic animal.

of interest. These systems provide little ability,

For many applications it is necessary to accurately control the time and pattern of transgene expression within an individual transgenic animal. For example, many attempts have been made to produce transgenic pigs which express increased levels of growth hormone (Vize et al., 1988, J. Cell Sci. 90:295-300;; Pinkert et al., 1990, Dom. Animal Endocrinol. 7:1-18). Elevated growth hormone levels 10 dramatically decrease the amount of body fat in pigs, and increase the animals overall feed efficiency. These effects would be beneficial, both to the consumer who could purchase a leaner, healthier product, and to the producer who can profit from having a more efficient animal. To date however, all 15 attempts to increase the level of growth hormone through production of transgenic pigs have also produced serious pathological conditions which greatly reduce the health of the animals. These pathologies 20 are the direct result of uncontrolled, constitutive expression of growth hormone, since many studies using exogenous hormone administration for short periods of time have not produced pathologies, while still benefiting feed efficiency and fat content. In this situation, a regulatory method to control onset and 25 level of expression from a growth hormone transgene would be extremely useful.

2.2. REPRESSOR-MEDIATED GENE CONTROL

DNA binding proteins with at least two functional sites. One site on the protein is used to bind DNA. The DNA binding site binds to a defined DNA sequence which is known as the operator site. Operator sites usually consist of palindromic sequences of 12 or more base pairs. A gene which is regulated by a repressor

must have at least one operator site located within its promoter/regulatory region. A second site on the repressor protein binds a specific ligand, usually a small macromolecule such as an amino acid, sugar, or antibiotic. When the ligand is bound to the repressor, it causes a conformational shift such that the affinity of the repressor for the operator sequence is greatly reduced. For this reason, the ligand is frequently referred to as the "inducer", since it causes the repressor to disassociate from the operator, thereby eliminating the repressor's effect and allowing expression of the gene.

Only the bacterial repressors Lacl, LexA and tetR have been shown to function in mammalian (Lacl and 15 LexA) or plant (tetR) tissue culture cells. The first report of utilizing bacterial repressors in eukaryotes was from Brent and Ptashne who showed that LexA could function in yeast (1984, Nature 312:612-615). Subsequently, both LexA and Lacl have been shown to 20 function in mammalian tissue culture systems (Smith et al., 1988, EMBO J. <u>7</u>:3975-3981). Of these repressors Lacl has been most extensively studied. For Lacl repression, single or multiple operator sites have been positioned in three major locations: (i) between 25 the transcription start site and the first codon of the mRNA; (ii) between the TATA-box sequence and the transcription start site; and (iii) between the TATAbox sequence and any more distal regulatory signal These studies reveal two predominant sequences. 30 results. First, operators located in all three positions were effective in rendering the modified promoter subject to Lacl repression. Second, the presence of multiple operator sequences allowed greater 1 vels of repression than did single operator 35

insertions. From these studies it appears the Lacl

repressor causes repression of mammalian promoters through two basic mechanisms. If the operators are located downstream of the transcription start site, Lacl appears to block expression by inhibiting mRNA elongation. That is to say, the LacI repressor blocks the progress of RNA polymerase by steric interference. When operator sequences are located in other positions, LacI seems to inhibit protein-protein interactions between the cellular factors normally involved in transcription initiation.

Gatz and Quail (1988, Proc. Natl. Acad. Sci. U.S.A. 85:1394-1397) have demonstrated tetR function in a plant protoplast culture system. Plant 15 protoplasts were transfected with a tetR gene expressed from a cauliflower mosaic virus (CAMV) promoter along with a CAT reporter gene, regulated by a modified CAMV promoter. In contrast to the results with Lacl, Gatz and Quail showed that tetR operators 20 positioned between the transcription start site and the first codon of the CAT mRNA were not responsive to tetR repression. Therefore the tetR protein does not appear to be able to block the procession of RNA polymerase. Effective repression by tetR was only 25 observed when the operator sequence was positioned such that the CAMV TATA-box element was flanked by the two 19bp palindromes of the tetR operator. With this modification, effective repression of the reporter gene, and induction with tetracycline could be 30 achieved. This suggests that repression by tetR specifically inhibits the initiation of transcription, in this case apparently by blocking the binding of the TATA-box binding factors.

Recently the tetR system has been shown to function in transgenic plants. Gatz et al. (1991, Mol. Gen. Genet. 227:229-237) have introduced their

original tetR responsive CAMV promoter, in which the operator sites flank the TATA-box into transgenic tobacco plants. Unexpectedly, this promoter, which exhibited very good regulation in tissue culture assays was not very effective in regulating gene expression in transgenic plants. Instead they found that effective repression and induction in transgenic plants occurred when the operator sites were positioned just downstream of the normal transcription start site.

3. SUMMARY OF THE INVENTION

The present invention relates to a tetracycline
repressor-mediated binary regulation system for the
control of gene expression in non-human transgenic
animals. It is based, at least in part, on the
discovery that in transgenic mice carrying two
transgenes, the first encoding bovine growth hormone
(bGH) under the control of a PEPCK promoter modified
to comprise the tetR operator sequence at the NheI
site, and the second encoding tetR repressor protein
under the control of an unmodified PEPCK promoter,
expression of bGH could be efficiently and selectively
induced by administering tetracycline to the
transgenic mice.

In particular embodiments, the present invention provides for (i) animal promoter elements modified to comprise a tetR operator sequence; (ii) nucleic acid molecules comprising a gene of interest under the control of such a modified promoter; (iii) non-human transgenic animals that carry a transgene under the control of said modified promoter and/or a transgene encoding the tetR repressor protein; and (iv) a method of selectively inducing the expression of a gene of interest in a non-human transgenic animal comprising

administering tetracycline to a non-human transgenic animal that carries a first transgene, which is the gene of interest under the control of a promoter modified to comprise a tetR operator sequence and a second transgene encoding the tetR repressor protein.

The present invention offers the advantage that, in the absence of tetracycline, expression of the gene of interest occurs at only very low levels due to efficient repression by tetR. In preferred, non-limiting embodiments of the invention, repression by tetR is further enhanced by utilizing a synthetic tetR gene which is devoid of splice signals and has optimized codon usage for mammalian cells.

Accordingly, the present invention allows tight control of gene expression in transgenic animals by withholding or administering tetracycline.

4. DESCRIPTION OF THE FIGURES

- Figure 1. A. Nucleotide sequence of tetR operator 20 as it occurs in Tn10, and in the oligonucleotides used to produce the modified PEPCK promoter elements. Bold face lettering represent the OP1 and OP2 tetR binding sites. The general purpose oligonucleotide is the sequence from $p\partial \partial 7$. 25 flanking EcoRI and AccI restriction sites used to excise this operator sequence are indicated. Additional restriction sites present in the plasmid, but not indicated here, which can be used to excise the operator include PstI, BamHI, 30 SpeI, Sbal, NotI, EagI, SacII, BstXI, and SacI on the 5' side and XhoI, ApaI and KpnI on the 3' The sequence of the PEPCK-TATA box operator is also indicated (see methods).
- 35 Figure 1. B. Nucleotide sequence of the $\partial \partial 7$

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operator. Lower case letters correspond to polylinker sequence. The 5' EcoRI and 3' AccI restriction sites used for producing the modified PEPCK promoters (Pck_A and Pck-N) are indicated. The 10 base pair linker beween OP1 and OP2 is underlined. Additional polylinker restriction sites available in pdd7 include PstI, BamHI, SpeI, XbaI, NotI, EagI, SacII, BstXI, and SacI on the 5' side and XhoI, ApaI and KpnI on the 3' side.

Figure 2. A representation of the three modified

PEPCK promoter elements. Construct 251 has the

\$\partial \partial 7\$ operator sequence integrated in the AccI site

of PEPCK, just 5' of the TATA-box control

element. Construct 252 has the \$\partial 7\$ operator

sequence incorporated into the NheI site of

PEPCK, just 3' of the TATA-box element.

Construct 261 incorporates the TATA-specific

operator sequence which is integrated between the

5' AccI site and the 3' NheI sites.

Figure 3. Structure of the modified PEPCK controlled bovine growth hormone genes. The Pck_AbGH and Pck_NbGH genes differ only in the site of operator insertion. For Pck_AbGH the operator is inserted at the AccI site 5' of the PEPCK TATA-box element. For Pck_NbGH the operator is inserted into the NheI site 3' of the TATA-box element (pPCK_NbGH has been deposited with the ATCC and assigned accession No:). In the Pck_TbGH gene, a TATA-box specific oligonucleotide was used, and this sequence was inserted between both the AccI and NheI sites.

A. Indicated the probe used for Sl hybridization.

35 Figure 4. S1 Nucleas protection assay to map the 5'

start site of bGH from the Pck N promoter. liver RNA (10µg) was hybridized to a 280 bp 5' labelled probe from the Pck NbGH gene in 40mM PIPES (Ph6.4), 1Mm EDTA, 400mM NaCl, 80% 5 formamide at 55° overnight. The probe spanned from the HinfI site in the 5' untranslated leader sequence of bGH to the PvuII site 5' of the TATA-17 box. The probe includes the tet-operator sequence of Pck N (see Figure 3). After 10 hybridization 300 μ l of ice cold digestion buffer (280mM NacL, 50Mm SODIUM ACETATE (Ph4.5), 4.5Mm ZnSO₄, 20µg/ml carrier DNA and 500 units S1 nuclease) was added and incubated at 37° for 30 The reaction as stopped by adding 80µl 15 of Stop Buffer (4M Ammonium acetate, 50mM EDTA and 50µg/ml tRNA), extracted with phenol/chloroform, precipitated with ethanol and analyzed on a 6% sequencing gel. The arrow indicates the protected fragment. Initiation of 20 bGH mRNA from the modified Pck_N promoter occurs approximately 20 bp 3' of the TATA-box. initiation site places the start of the message just prior to the first tetR binding site. result indicates that the bGH mRNA starts from a 25 single cap site, and suggests that tetR repression is due to a block in transcription initiation. Furthermore, unrepressed bGH expression appears to be due to limited tetR expression. 30

- Figure 5. Nucleotide sequence of the tetR repressor protein gene.
- Figure 6. Alterative, nonlimiting promoters of interest. Asterisks indicate sites at which tetR operator sequence may be inserted.
- Figure 7. Northern blot analysis of bGH mRNA in liver

of F1 generation animals.

- Figure 8. Northern blot analysis of bGH mRNA expression in four transgenic lines.
- 5 Figure 9A. Tissue specificity of bGH expression in Line 10-2 in the presence of 50 µg/ml tetracycline. Northern blot analysis of bGH induction in a variety of tissues. Only the liver and kidney show significant expression.
- Pigure 9B. Tetracycline induction of bGH in Line 10-2.

 Both liver and kidney, which are the only sites
 for bGH expression in Figure 9A, also show
 tetracycline dependent bGH expression.

Figure 10. 345 Repressor Construct.

- 15 Figure 11. Induction of bGH expression in Construct
 345 Offspring. Northern blot analysis of liver
 RNA from F1 animals containing the 345 construct.
 Only animals from line 14 exhibit tetracycline
 dependent bGH expression.
- Figure 12. Expression and alternative processing of tetR transgene. A RNase protection probe which extends from the Nrul site of tetR 3' to the end of the gene was used. This probe includes only tetR coding sequences and should give a fully protected fragment of approximately 400 base pairs. A protected fragment of approximately 220-260 base pairs is observed, which is far smaller then predicted.
- Figure 13. 5' Structure of tetR mRNA. Liver RNA was treated with reverse transcriptase and amplified by PCR. The RNA was amplified using two different pairs of primers. The first primer pair (TZ-1 and TZ-4) should produce a 619 base pair product. The second primer pair (TZ03 and TZ04) should produce a 498 base pair product. The sequence of the primers ar:

TZ-1: 5 CCGCATATGATCAATTCAAGGCCGAATAAG3 TZ-3: 5'CTTTAGCGACTTGATGCTCTTGATCTTCCA3' TZ-4: 5'AATTCGCCAGCCATGCCAAAAAAGAAGAGG3' The TZ-4 primer is common to both primer pairs 5 and is the 5' primer which encompasses the start codon of the tetR and mRNA. Primer TZ-1 and TZ-3 are two different 3' primers both of which are in the tetR coding region. When amplified, these primer pairs produced smaller then expected 10 products (approx. 215bp vs. 619bp for TZ-4 and TZ-1, and approx. 94bp vs. 498bp for TZ-4 and TZ-3). The products of this reaction were cloned and sequenced. Sequencing revealed the presence of an unexpected intron which spanned from near 15 the Xbal site at the start of tetR to a splice acceptor just 8 base pairs 5' of the TZ-3 primer. Figure 14. Composition analysis of Wild Type Tn10 tetR gene. The Tn10 tetR coding sequence was analyzed on a desktop computer using Mac Vector 20 The figure shows a diagram of the tetR coding region with the plus strand splice doner (D) and splice acceptor (A) signal sequences indicated. For reference the location of the XbaI restriction is also indicated. The first 25 graph depicts the percentage of G and C bases in the coding region of tetR. There are several domains of very low GC content. The bottom graph is an analysis of codon bias. The dark line is a comparison of the tetR codon usage to a mouse 30 codon bias table. Values lower than 1.0 are indicative of sequences which may translate poorly. For reference, a comparison of tetR to a Tobacco codon bias table is included (light

line). In transgenic tobacco, the tetR

regulation system functions very efficiently,

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suggesting that for this gene, codon bias may be an important factor for efficient expression.

Figure 15. Synthetic tetR Component Sequences. The components of the synthetic tetR gene were synthesized by Midland Laboratories as four overlapping double stranded DNA cassettes. The sequence of these cassettes are shown. Each cassette was blunt cloned into the Hinc2 site of pUC19 and sequenced to verify authenticity. The resulting plasmids pLT1, pLT2, pLT3 and pLT5 can be used as the source material to assemble the entire synthetic tetR coding sequence since each contains an overlapping unique restriction site (bold face) through which they can be joined.

Figure 16. Sequence of Synthetic tetR gene.

Figure 17. Composition analysis of synthetic tetR.

These graphs were produced using the same software described in Figure 15. The figure depicts the structure of the synthetic tetR gene,

now devoid of splice donor signal sequences, with only a single splice acceptor signal remaining (A). This is not the splice acceptor which was active in the 345 construct. The percentage of G and C bases has been significantly improved,

while the frequency of CpG base pairs has been kept to a minimum. A CpG base pair is frequently the site for DNA methylation, which can negatively effect the expression of a gene. The

codon bias of the synthetic tetR gene is also vastly improved. The graph depicts the results when the synthetic tetR coding sequence is compared to the same mouse codon bias table used previously.

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5. DETAILED DESCRIPTION OF THE INVENTION

For purposes of clarity of description, and not by way of limitation, the detailed description of the invention is divided into the following subsections:

- (i) the tetR operator;
- (ii) modified promoters containing the tetR
 operator; and
- (iii) utility of the invention.

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5.1. THE TETR OPERATOR

In order to practice the instant invention, the tetR operator sequence is inserted into a suitable animal promoter sequence in order to render that promoter subject to control by tetR repressor protein. A diagram of the tetR operator sequence is depicted in Figure 1.

It may be convenient to clone the tetR operator into a vector, such as a plasmid or a phage, to facilitate its propagation. Cloned operator sequence may then be rendered available for insertion into a promoter of interest, as set forth in Section 5.2., infra.

In a particular, nonlimiting embodiment of the invention, tetR operator sequence may be cloned as follows: Four oligonucleotides, which when annealed produce the two 19bp OP1 and OP2 palindromic sequences of the tetR operator may be synthesized; the sequences of said oligonucleotides are as follows:

- 30 X-1. 5'ACTCTATCATTGATAGAGT3'
 - X-2. 5'ACTCTATCAATGATAGAGT3'
 - X-3. 5'TCCCTATCAGTGATAGAGA3'
 - X-4. 5'TCTCTATCACTGATAGGGA3'

Oligonucleotides X-1 and X-2 are complementary and,

when annealed, form the OP1 operator. Similarly, oligonucleotid s X-3 and X-4, when annealed, produce

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the OP2 operator site. The OP1 oligonucleotides may then be directly cloned into the EcoRV site of the Bluescript (Stratagene) polylinker to form plasmid X. OP2 oligonucleotides may then be cloned into a Mung bean nuclease blunted ClaI site of plasmid X to form plasmid Y. The resulting tetR operator may then be propagated and then excised from plasmid Y as an EcoRI, AccI fragment which may be end-filled with T4 polymerase and gel purified.

It is preferable that the separation between OP1 and OP2 is about 10-11 bp.

Analogous methods may be used to insert the tetR operator site into other suitable vectors.

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5.2. MODIFIED PROMOTERS CONTAINING THE tetr OPERATOR

According to the invention, the tetR operator may be inserted into a suitable animal promoter so as to render that promoter subject to repression by tetR repressor protein. Any animal promoter maybe used; strategies for promoter selection are set forth in Section 5.3., infra.

In preferred embodiments of the invention, the tetR operator sequence is positioned 3' to the TATA-box sequence. A nonlimiting list of promoters which may be used according to the invention is set forth in Figure 6, together with the proximal portion of the promoter in the vicinity of the TATA-box, which is underlined.

In a specific, nonlimiting embodiment of the invention, the tetR operator site may be inserted into the NheI site of the PEPCK promoter (Wynshaw-Boris et al., 1984, J. Biol. Chem. <u>259</u>:12161-12169). A diagram of the PEPCK promoter containing the tetR operator sequence of the NheI site is presented in Figure 2.

For insertion of the operator sequence, the PEPCK promoter may be cut with NheI and end-filled with T4 polymerase; tetR operator, prepared as set forth in Section 5.1., supra, may then be blunt-ligated into place.

5.3. UTILITY OF THE INVENTION

5.3.1. STRATEGY

non-human transgenic animal that comprises two transgenes. The first transgene, termed "A," is a gene of interest, the expression of which is desirably controlled. Virtually any gene of interest may be used, including, but not limited to, growth hormone, hemoglobin, low density lipoprotein receptor, insulin, genes set forth in Table I, etc.

TABLE 1

Other Genes Of Interest

Gene	Diagram / Afficiati
Gene	Disease/Affect
ADA Adenosine deaminase	Immuno-deficiency
TNF Tumor necrosis factor	Anti-cancer
IL-2 Interleukin-2	Anti-cancer
LDL low density	hypercholesterolemia
Factor IX	hemophelia
Factor VIII	hemophelia
eta-glucosidase	Gauchers disease
CFTR Cystic fibrosis	Cystic fibrosis
transmembrane regulator	
HPRT Hypoxanthine-guanine	Lesch-Nyhan syndrome
phosphoribosyltransferase	
UDP-glucuronyl transferase	Crigler-Najjar syndrome
Growth Hormone receptor	Growth
Insulin-like growth factor	Growth
Growth hormone releasing	Growth
factor	

The expression of gene "A" is under the transcriptional control of promoter "B". Promoter B comprises a tetR operator sequence, as discussed supra. Promoter B desirably defines the time and tissue window in which the transgene may be induced; for example, promoter A may be a tissue specific promoter such as the PEPCK promoter (which is expressed selectively in liver and becomes active shortly prior to birth). The second transgene encodes the tetR repressor, the sequence of which is set forth in Figure 5.

Analysis of the Tn10 tetR coding sequence indicates that the codon usage for this gene is poorly suited for expression in mammalian cells (FIG. 15). 15 To optimize tetR expression in mammalian cells a new tetR repressor gene was designed (See, Section 7, infra), which may be utilized in alternative embodiments of the invention. The synthetic tetR gene (syn-tetR) is designed to encode exactly the same 20 protein product as the bacterial Tn10 tetR gene but optimizes codon usage for mammalian cells. percentage of G and C bases has been significantly improved, while the frequency of CpG base pairs has been minimized. A CpG base pair is frequently the 25 site for DNA methylation which can negatively affect the expression of a gene. In addition, the syn-tetR gene is devoid of any splice signals, decreasing the likelihood of aberrant splicing of the RNA which may result in production of a non-functional message. sequence of the synthetic tetR gene is depicted in Figure 16. Plasmids comprising these sequences may be constructed using plasmids pLT-1, pLT-2, pLT-3 and pLT-5 (deposited with the American Type, Culture Collection (ATCC) and assigned accession numbers 35

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_, ____, and ____, as described in Section 7, infra.

In further embodiments, the present invention 5 provides for additional synthetic tetR genes from which one or more splice sites have been deleted or for which codon usage has been further optimized.

The present invention covers synthetic tetR genes having the sequence set forth in Figure 16 and for functionally equivalent variants of that sequence.

In specific, non-limiting embodiments of the invention, a nuclear localization signal may be added to a natural or synthetic tetR gene to facilitate its expression (See, Section 7, infra).

Expression of tetR is controlled by promoter "C". While it is preferable that promoter C be the same as promoter B except that promoter C does not contain a tetR operator sequence, any promoter which provides expression of tetR so as to repress expression of gene "A" during the period when it is desirable to repress expression of "A" may be used.

For example, and not by way of limitation, a transgenic animal may be produced which carries a first transgene which is bovine growth hormone under 25 the control of a PEPCK promoter modified to contain a tetR operator sequence at the NheI site and a second transgene which is tetR repressor protein under the control of an unmodified PEPCK promoter; see Section 6, <u>infra</u>. The pPCK NbGH construct has been deposited with the ATCC and assigned accession number

5.3.2. TRANSGENIC ANIMALS OF THE INVENTION

The binary repressor system of the invention may be used to control gene expression in any non-human 35 transgenic animal, including, but not limited to, transgenic mice, pigs, goats, cows, rabbits, sheep,

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etc. The present invention provides for such nonhuman transgenic animals carring as transgenes nucleic acid constructs described herein, including natural or synthetic tetR repressor proteins and operator sequences.

Transgenes may be introduced by microinjection, transfection, transduction, electroporation, cell gun, embryonic stem cell fusion, or any other method known in the art. The transgenes of the invention may be co-introduced into a single animal or may be introduced into two individual animals that are subsequently mated to produce doubly transgenic offspring.

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mother.

For example, for the production of transgenic 15 mice, the following general protocol may be used. Male and female mice are mated at midnight. hours later, the female may be sacrificed and the fertilized eggs may be removed from the uterine tubes. 20 Foreign DNA may then be microinjected (100-1000 molecules per egg) into a pronucleus. Shortly thereafter, fusion of the pronuclei (a pronucleus or the male pronucleus) occurs, and, in some cases, foreign DNA inserts into (usually) one chromosome of 25 the fertilized egg or zygote. The zygote may then be implanted into a pseudo-pregnant female mouse (previously mated with a vasectomized male) where the embryo develops for the full gestation period of 20-21 The surrogate mother then delivers the mice and by four weeks transgenic pups may be weaned from the 30

According to another embodiment of the invention, a transgenic pig may be produced, briefly, as follows. Estrus may be synchronized in sexually mature gilts (>7 months of age) by feeding an orally active progestogen (e.g. allyl trenbolone, AT: 15mg/gilt/day)

gilts may be given an intramuscular injection of prostaglandin F_{2a} (Lutalyse: 10mg/injection) at 0800 and 1600 hours. Twenty-four hours after the last day of AT consumption all donor gilts may be administered a single intramuscular injection of pregnant mare serum gonadotrophin (1500 U). Human chorionic gonadotrophin (750 IU) may be administered to all donors at 80 hours after pregnant mare serum gonadotrophin.

Following AT withdrawal, donor and recipient gilts may be checked twice daily for signs of estrus using a mature boar. Donors which exhibited estrus within 36 hours following human chorionic gonadotrophin administration may be bred at 12 and 24 hours after the onset of estrus using artificial and natural (respectively) insemination.

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Between 59 and 66 hours after the administration of HCG one- and two-cell ova may be surgically 20 recovered from bred donors using the following procedure. General anesthesia may be induced by administering 0.5 mg of acepromazine/kg of bodyweight and 1.3 mg of ketamine/kg via a peripheral ear vein. Following anesthetization, the reproductive tract may 25 be exteriorized following a mid-ventral laparotomy. A drawn glass cannula (O.D. 5 mm, length 8 cm) may be inserted into the ostium of the oviduct and anchored to the infundibulum using a single silk (2-0) suture. Ova may then be flushed in retrograde fashion by 30 inserting a 20g needle into the lumen of the oviduct 2 cm anterior to the uterotubal junction. Sterile Dulbecco's phosphate buffered saline (PBS) supplemented with 0.4% bovine serum albumin (BSA) may be infused into the oviduct and flushed toward the 35 glass cannula. The medium may b collected into

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sterile 17 x 100 mm polystyrene tubes. Flushings may be transferr d to 10 x 60 mm petri dishes and searched at a lower power (50x) using a Wild M3 stereomicroscope. All one- and two- cell ova may be washed twice in Brinster's Modified Ova Culture -3 medium (BMOC -3) supplemented with 1.5% BSA and transferred to 50 μ l drops of BMOC-3 medium under oil. Ova may be stored at 38°C under a 90% Nz, 5% Oz, 5% Co2 atmosphere until microinjection is performed. One and two-cell ova may be placed in an Eppendorf tube (15 ova per tube) containing 1 ml HEPES medium supplemented wit 1.5% BSA and centrifuged for 6 minutes at 14,000g in order to visualize pronuclei in one-cell and nuclei in two-cell ova. Ova may then be transferred to a 5-10µl drop of HEPES medium under oil on a depression slide. Microinjection may be performed using a Laborlux microscope with Nomarski optics and two Leitz micromanipulators. molecules of construct DNA (linearized at a concentration of about 1ng/µl of Tris-EDTA buffer) may be injected into one pronucleus in one-cell ova or both nuclei in two-cell ova. Microinjected ova may be returned to microdrops of BMOC-3 medium under oil and 25 maintained at 38°C under a 90% N₂, 5% CO₂, 5% O₂ atmosphere prior to their transfer to suitable recipients. Ova may preferably be transferred within 10 hours of recovery. Only recipients which exhibit estrus on the same day or 24 hours later than the

30 donors may preferably be utilized for embryo transfer. Recipients may be anesthetized as described supra. Following exteriorization of one oviduct, at least 30 injected one- and/or two-cell ova and 4-6 control ova may be transferred in the following manner. tubing from a 21g x 3/4 butterfly infusion set may be 35

connected to a lcc syringe. The ova and one to two

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mls of BMOC-3 medium may be aspirated into the tubing. The tubing may then be f d through the ostium of the oviduct until the tip reaches the lower third or isthmus of the oviduct. The ova may be subs quently expelled as the tubing is slowly withdrawn. exposed portion of the reproductive tract may be bathed in a sterile 10% glycerol - 0.9% saline solution and returned to the body cavity. connective tissue encompassing the linea alba, the 10 fat, and the skin may be sutured as three separate An uninterrupted Halstead stitch may be used to close the linea alba. The fat and skin may be closed using a simple continuous and mattress stitch, respectively. A topical antibacterial agent (e.g. 15 Furazolidone) may then be administered to the incision Recipients may be penned in groups of about four and fed 1.8 kg of a standard 16% crude protein corn-soybean pelleted ration. Beginning on day 18 (day 0 = onset of estrus), all recipients may be 20 checked daily for signs of estrus using a mature boar. On day 35, pregnancy detection may be performed using ultrasound. On day 107 of gestation recipients may be transferred to the farrowing suite. In order to ensure attendance at farrowing time, farrowing may be 25 induced by the administration of prostaglandin F_{2a} (10) mg/injection) at 0800 and 1400 hours on day 112 of gestation. In all cases, recipients may be expected to farrow with 34 hours following PGF 2a administration. 30

As used herein, the term "transgenic animal" refers to animals that carry a transgene in at least some of their somatic cells, and preferably in at least some of their germ cells.

5.3.3. INDUCTION

Induction of expression of the gene of interest in transgenic animals of the invention may be achieved by administering, to the animal, a compound that binds to tetR so that tetR repressor function is inhibited. Examples of such compounds include tetracycline and tetracycline-like compounds, including, but not limited to, apicycline, chlortetracycline, clomocycline, demeclocyline, guamecycline, lymecycline, meclocycline, methacycline, minocycline, oxytetracycline, penimepicycline, pipacycline, rolitetracycline, sancycline, and senociclin.

Administration of the inducer can be through direct injection, water, feed, aerosol, or topical 15 application. The choice of method will depend on the promoters used and the specific application of the transgenic animals. For example, injection, water and feed would provide inducer to all of the animals tissues. In our case, administration through water or 20 feed would be the preferred method to control growth hormone expression in transgenic pigs. Aerosol spray could be used to attain high antibiotic concentrations in the lung. This may be appropriate for example in a 25 cystic fibrosis or emphysema model. Topical application to the skin is also possible and could be used in models of acne, hair loss, wound healing or viral infection.

Induction of the gene of interest is accomplished by administering an effective amount of inducer, as described above. An effective amount of inducer may be construed to mean that amount which increases expression of the gene of interest by at least about 50 percent. As the LD₅₀ for tetracycline HCl in rats is about 6643 mg/kg and the therapeutic dose is between about 25-50 mg/kg, an effective dose of

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tetracyclin, as inducer, is between about 5-50 mg/kg and preferably betw en ab ut 5-15 mg/kg.

6. EXAMPLE: TETRACYCLINE REPRESSOR-MEDIATED BINARY REGULATION SYSTEM FOR CONTROL OF BOVINE GROWTH HORMONE EXPRESSION IN TRANSGENIC MICE

6.1. MATERIALS AND METHODS

6.1.1. CONSTRUCTION OF PLASMIDS

Plasmid pdd7 contains a functional tetR operator site cloned within a Bluescript (Stratagene) polylinker. This plasmid is useful for propagating the operator sequence, and as a source of operator sites for insertion into the PEPCK promoter or any other promoter element. The pdd7 plasmid was made as follows. Four oligonucleotides, which when annealed produce the two 19bp OP1 and OP2 palindromic sequences of the tetR operator were synthesized. The sequences of each oligonucleotide is listed below.

X-1.5' ACTCTATCATTGATAGAGT 3'

X-2.5' ACTCTATCAATGATAGAGT 3'

X-3.5' TCCCTATCAGTGATAGAGA 3'

X-4.5' TCTCTATCACTGATAGGGA 3'

Oligonucleotides X-1 and X-2 are complementary and when annealed form the OP1 operator. Similarly oligonucleotides X-3 and X-4 produce the OP2 operator site. The OP1 oligonucleotides were directly cloned into the EcoRV site of the Bluescript polylinker. The resulting plasmid pSOPI was sequenced to verify the integrity of the insert. OP2 oligonucleotides were subsequently cloned into a Mung bean nuclease blunted Clal site of pSOPI to produce $p\partial\partial T$. Due to a cloning artifact produced by the Mung bean nuclease, the operator in $p\partial\partial T$ consists of the two 19bp OP1 and OP2 sequences separated by linker of only 10 base pairs. This difference does not effect tetR binding. The

sequence of the pdd7 operator sit is shown in Figure 1B. The 55 base pair t tR operator was excised from pdd7 as an EcoRl, AccI fragment, end filled with T4 polymerase, and gel purified. This fragment was subsequently used to produce the modified PEPCK promoters Pck N and Pck A.

Plasmids Pck A and Pck N were produced by inserting the 55bp tetR operator into the unique AccI and NheI sites (respectively) of the PEPCK promoter 10 (pPCK_NbGH has been deposited with ATTC and assigned For both plasmids the promoter was accession No:) . cut with the appropriate restriction enzyme, end filled with T4 polymerase and the tetR operator blunt 15 ligated into place. A third modified PEPCK promoter, Pck T was produced in which the OP1 and OP2 operator sequences were positioned to flank the PEPCK TATA-box To produce Pck_T a new oligonucleotide element. (5'ACTCTATCATTGATAGAGTTACTAT

TTAAATCCCTATCAGTGATAGAGA3') was produced. This oligonucleotide was kinased with T4 polynucleotide kinase and annealed to kinased X-2 and X-4 which are complementary to the first and last 19bp. The complete double stranded 49bp operator was produced by filling in the 11bp linker region, which includes the PEPCK TATA-box element, with Klenow. The final product was then blunt cloned into an AccI, NheI cut PEPCK promoter. All three modified promoters were sequenced to verify the inserts. Figure 2 depicts the structure of these promoters.

6.1.2. REPRESSOR CONSTRUCT

Plasmid pBI501 contains a 701 bp HincII fragment from E. coli Tn10, cloned into the HincII site of pUC8. The HincII insert contains the entire tetR coding sequence along with 21bp of 5' and 55bp of 3'

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untranslated DNA. This insert was excised from the parent plasmid and subcloned into a plasmid with a more suitable polylinker to produce pSTET7. To this plasmid a 870bp XhoI, BamHI fragment derived from pMSG (Pharmacia), containing the SV40 small-T intron and polyadenylation signal sequences was inserted at the HindII site 3' of the tetR coding region to produce pSTetRSv. Finally an unmodified 610bp PEPCK promoter was inserted at the EcoRl site of pSTETRSv to produce pPck_tetRSv. The PEPCK promoter is identical to the promoter used to produce pPck_A, pPck_N, and pPck_T except that it does not contain a tetR operator site. This PEPCK promoter has been previously used in transgenic animals and is known to target gene expression specifically to the liver.

6.1.3. GROWTH HORMONE GENES

Plasmid pGH-SAF107 contains a 2.2kb BamHI, EcoRI genomic fragment of the bovine growth hormone (bGH) 20 gene, blunt ligated into an EcoRV site. vector each of the modified PEPCK promoters was added by blunt ligating the promoter into the BamHl site of The structure of the resulting plasmids pGH-SAF107. is depicted in Figure 3. Plasmid pPCK_NbGH was 25 deposited with the ATCC and assigned accession number For production of transgenic animals, each of the PEPCK-bGH genes was excised from the vector using Xhol and Sacl, gel fractionated and purified using an Elutip column. 30

6.1.4. TRANSGENIC MICE

Transgenic mice were made which contain both the Pck_tetRSv gene and one of the modified PEPCK promoters controlling bGH. Table 2 lists the number

of eggs injected, offspring produced and number of transgenics derived for ach construct.

TABLE 2

Construct	Eggs injected	Eggs transferred	Live Born	Transgenic
Pck_AbGH + Pck_tetRSv (251)	233	194	40	14 (0.35)
Pck-NbGH + Pck_tetRSv (252)	268	208	30	9(0.3)
Pck_TbGH + Pck-tetRSv (261)	227	197	25	5(0.2)

6.2. RESULTS AND DISCUSSION

Once the transgenic founder animals were identified, they were weighed each week. Table 3 lists the mean weights of each group of transgenic animal at 11 weeks of age.

TABLE 3

Construct	Sex	Weight
Pck_AbGH + Pck_tetRSv(9)	male	36.122(12.251)
Pck_AbGH + Pck_tetRSv(4)	female	29.125(7.861)
Pck_NbGH + Pck_tetRSv(5)	male	34.840(14.745)
Pck_NbGH + Pck_tetRSv(4)	female	28.125(10.958)
Pck_TbGH + Pck_tetRSv(3)	male	36.267(11.402)
Pck_TbGH + Pck_tetRSv(2)	female	27.300(5.798)
NON-TRANSGENIC(6)	male	29.583(2.395)
NON-TRANSGENIC(6)	female	23.117(1.863)

As expected for each co-injection, large animals, obviously expressing elevated levels of bGH, were observed as were animals of normal stature.

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At 10 weeks of age, a sampling of transgenic female founders containing the A+T and N+T were tested for induction of bGH in the serum using a radio-immune assay, after a single IP injection of 60 mg/kg tetracycline-HCl. The purpose of this experiment was simply to determine which if either of these two modified promoters was responsive to repression by tetR. The results are summarized in Table 4.

TABLE 4

Construct	Animal	Weight	Basal	12 hours	36 hours
249	2-5 female	21.1	0.00	0.00	0.00
250	6-6 female	42.9	4.6 <u>+</u> 0.033	3.4 <u>+</u> 0.062	4.9 <u>+</u> 0.072
251	6-6 female	19.3	0.00	0.00	0.00
251	10-5 female	25.1	0.20 <u>+</u> 0.008	0.19 <u>+</u> 0.001	0.21 <u>+</u> 0.038
252	5-2 female	38.7	0.59 <u>+</u> 0.107	0.64 <u>+</u> 0.044	1.12 <u>+</u> 0.207
252	5-3 female	20.0	0.00	0.00	0.00
252	10-2	19.2	0.00	0.00	0.00

No induction of bGH was observed in animals that lack the Pck tetRSV gene (construct 250) or in animals with 25 both the Pck_AbGH + Pck-tetRSv genes (construct 251). An approximate two fold increase in serum bGH levels was however detected in the 5-2 female which contains the Pck-NbGH + Pck_tetRSV genes. The remainder of the animals had undetectable levels of bGH expression, due in part to the relatively low sensitivity of this assay. For example the 10-2 female (construct 252) shows no detectable bGH in the serum, but subsequent experiments on her offspring indicate that this line of animals does express bGH mRNA in a tetracycline dependent manner. This initial data, suggested that

th Pck_N promoter was being r gulated by tetR at 1 ast to a limited extent.

sensitivity of the assay and to test the responsiveness of the Pck_T promoter, offspring of founder mice from each co-injection were produced. The transgenic progeny were then raised in the presence or absence of tetracycline medicated water (500μg/ml) for 4 weeks, prior to analysis of bGH mRNA expression levels in the liver, the predominant site of PEPCK expression. Northern blot hybridization analysis of these animals (Figure 7) demonstrated again, that animals with the Pck_NbGH gene were responsive to repression by tetR and that the other two modified promoters exhibited no signs of tetR dependent regulation.

We attempted to breed all of the remaining founders containing the Pck-NgGH + Pck tetRSv genes to analyze their offspring in a similar manner (Figure 20 8). Of the 5 founders which produced offspring, 2 did not express bGH under any conditions, and from the remaining 3 one segregated two different integration sites allowing us to establish a total of 4 lines. All 4 lines exhibited tetracycline dependent bGH 25 expression as assayed by Northern blot hybridization. The efficiency of tetR repression appeared to be inversely correlated with the level of expression. For example 9-5 animals have the highest level of bGH expression, show an obvious increase in body size, and 30 exhibit only marginal tetR repression. In contrast 9-4Lc and 10-2 animals exhibit lower levels of tetracycline induced bGH expression, are of normal stature and appear to be efficiently regulated by t tR. 35

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An S1 nucl ase protection assay was performed to identify the start site of transcription of bGH mRNA. As shown in Figure 4, there was only one start site identified regardless of the presence or absence of tetR repressor binding. This start site was located approximately 20 bp downstream from the TATA-box. At this location, the message is initiating within the dd7 operator sequence, just 3 or 4 base pairs 5' of the first tetR binding site.

7. EXAMPLE: OPTIMIZATION OF tetR CODING SEQUENCE

The use of the wild type Tn10 tetR gene in conjunction with the 252 construct indicates that the TetR system can function in transgenic animals and that in some cases, for instance in the 10-2 transgenic animals, the level of regulation can be very high (FIGS. 9A and 9B). However, in other instances the efficiency of repression is not always complete, leading to a significant basal level of bGH expression. This failure to repress may be due to low level expression of tetR. To optimize the expression of tetR repressor, a synthetic tetR gene was generated which was devoid of splice signals and had optimized codon usage for mammalian cells.

7.1 MATERIALS AND METHODS

7.1.1. TISSUE SPECIFICITY AND TETRACYCLINE INTRODUCTION OF bGH IN LINE 10-2

For all Northern blots $10\mu g$ of whole RNA was electrophoreses through a 1% agarose gel containing 3% formaldehyde using standard techniques. To detect bGH mRNA a random primed, radioactive bGH cDNA probe was used. All conditions for hybridization and washing of

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filters were done in accordance with standard techniques of mol cular biology.

7.1.2. EXPRESSION AND ALTERNATIVE PROCESSING OF THE tetr TRANSGENE

A RNase protection probe which extended from the NruI site of tetR 3' to the end of the gene was used. This probe includes only tetR coding sequences and should give a fully protected fragment of approximately 400 base pair. When hybridized to 150µg of liver RNA (500,000 cpm of probe in a 30µl hybridization consisting of 80% formamide, 40mM PIPES pH 6.4, 400mM NaOAc, and 1mM EDTA), and digested with RNase one (Promega) for 30 minutes at 37° as recommended by the manufacturer, a protected fragment of approximately 221-260 base pairs is observed, far smaller than predicted.

7.1.3. 5' STRUCTURE OF tetr mRNA

Liver RNA was treated with reverse transcriptase and amplified by PCR using the manufacturers recommended conditions (Pharmacia). The RNA was amplified using two different pairs of primers. The first primer pair (TZ-1 and TZ-4) should produce a 619 base pair product. The second primer pair (TZ-3 and TZ-4) should produce a 498 base pair product. The sequence of the primers are:

- TZ-1: 5'CCGCATATGATCAATTCAAGGCCGAATAAG3'
- 30 TZ-3: 5'CTTTAGCGACTTGATGCTCTTGATCTTCCA3'
 - TZ-4: 5'AATTCGCCAGCCATGCCAAAAAAGAAGAGGG3'

The TZ-4 primer is common to both primer pairs and is the 5' primer which encompasses the start codon of the tetR mRNA. Primer TZ-1 and TZ-3 are two different 3' primers both of which are in the t tR coding r gion. When amplified, these primer pairs

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produce small r than expected products (approx. 215bp vs. 619bp for TZ-4 and TZ-1, and approx. 94bp vs. 498bp for TZ-4 and TZ-3). The products of this reaction were cloned and sequenced. The sequence revealed the presence of an unexpected intron which spanned from near the Xbal site at the start of tetR to a splice acceptor just 8 base pairs 5' of the TZ-3 primer.

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7.1.4. 345 REPRESSOR CONSTRUCT

In an embodiment of the invention, any nuclear localization signal may be added to a natural or synthetic tetR gene to facilitate its expression. For example, complementary oligonucleotides which encode a nuclear localization signal sequence were synthesized (Oligos etc.) and added in frame to the tetR coding sequences of pSTETR107 at the EcoR1 and Xbal restriction sites to produce pNTETR. Oligonucleotide sequences are:

(GB1) 5'AATTCGCCAGCCATGCCAAAAAAGAAGAGGAAGGTAT3' and (GB2) 5'CTAGATACCTTCCTCTTCTTTTTTGGCATGGCTGGC3'. When annealed these oligonucleotides have a 5' EcoR1 and 3' Xbal compatible overhangs. These

oligonucleotides fuse the amino acid sequence Met Pro Lys Lys Lys Arg, Lys Val, to the third amino acid (Arg) of wild type tetR.

Two complementary 51 base pair oligonucleotides which start the 5' cap site of bGH and extend to the first exon were synthesized (Oligos etc.). Sequence for the oligonucleotides are (5b-1): 5'GATCCCAGGACCCAGTTCACCAGACGACTCAGGGTCCTGTGGACAGCT CAG3'

and (5b-2):

35 5 AATTCTGAGCTGTCCACAGGACCCTGAGTCGTCTGGTGAACTGGGTCC

TGG3'. When annealed these oligonucleotides have 5'
BamH1 and 3'EcoR1 compatible overhands. The
oligonucleotides for the 5' leader sequence of bGH
were cloned into a BamH1, EcoR1 cut plasmid to produce
p5'GH.

The nuclear localization modified tetR coding sequence was isolated by gel purification after restriction digestion of pNTETR using EcoR1 and Hind III. This fragment was then inserted into p5'GH at the EcoR1 and Hind III sites to product p5'GHTR.

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To add the remainder of the bGH genomic sequence an intermediate modification of p5'GHTR was first This modification consisted of adding a Hind III - Pst1 linker to the Hind III site of p5'GHTR 15 to product pGTO. The sequence of the oligonucleotides which comprise this linker are: (CC-1) 5'AGCTTCTGCAG3' and (CC-2) 5'AGCTCTGCAGA3'. remaining bGH genomic sequences were added in two 20 steps. First the Pst1 Sac2 fragment that begins in the first exon of bGH and ends in the third intron was excised from pSGH107. Similarly, the insert of pGTO which contains the 5' untranslated leader of bGH and the nuclear localization modified tetR was excised 25 using BamH1 and Pst1. These two gel purified fragments was then cloned into a BamH1 Sac2 cut vector to produce pGTG. Finally, the remainder of the bGH gene from the Sac2 site in the third intron to the end of the gene, was added to pGTG by cutting pGTG with Sac2 and adding the Sac2 fragment from pSGH106 to 30 produce pNTETR-GH.

Plasmid pNTETR-GH was digested with BamH1 to excise the NTETR-GH gene. The fragment was cloned into the BamH1 site of pPCK 305 to produce the final plasmid pPCK-GHNTET. To produce transgenic mice, the PEPCK-GHTET gene was excised from the plasmid using

Sall and Sacl. This fragment was gel purified and coinjected with the PCK-NbGH gene previously described to generate transgenic mice.

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SYNTHETIC tetR COMPONENT SEQUENCES

The components of the synthetic tetR gene were synthesized by Midland Laboratories as four overlapping double stranded DNA cassettes. sequence of these cassettes are shown in Figure 15. Each cassette was blunt cloned into the Hinc2 site of pUC19 and sequenced to verify authenticity. resulting plasmids pLT1, pLT2, pLT3 and pLT5 can be used as the source material to assemble the entire synthetic tetR coding sequence since each contains an overlapping unique restriction site (bold face) through which they can be joined (pLT-1, pLT-2, pLT-3 and pLT-5 have been deposited with ATCC and have been assigned accession numbers ____, ___, and ____ respectively). There are many possible ways by which 20 these cassettes can be joined. By way of an example, the inserts of plasmid pLT1 and pLT2 can be excised using EcoR1 and Nsi1. The inserts can then be combined by cloning these two fragments into an EcoR1 vector. This procedure will assemble the 5' half of the gene, using the overlapping Nsil restriction site to join the pieces. Similarly, the 3' half of the gene can be assembled from pLT3 and pLT5 by cutting with EcoR1 and Sph1 (pLT3) and Sph1 and Hind III (pLT5) to release the inserts. These inserts can then be joined at the overlapping Sph1 site by cloning the fragments into an EcoR1, Hind III cut vector. Finally, the entire coding region can be put together using the overlapping restriction site ApaL1. 35 would result in a vector with the synthetic tetR

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coding sequence, as depicted in Figure 16, cloned into a plasmid as an EcoR1 Hind III fragment.

7.1.6. COMPOSITIONAL ANALYSIS OF WILD TYPE Tn10 tetR GENE

The Tn10 tetR coding sequence was analyzed on a desktop computer using Mac Vector software. Figure 14 shows a diagram of the tetR coding region with all of the plus strand splice doner (D) and splice acceptor 10 (A) signal sequences indicated. For reference the location of the Xbal restriction is also indicated. The first graph depicts the percentage of G and C bases in the coding region of tetR. There are several domains of very low GC content. The bottom graph is 15 an analysis of codon bias. The dark line is a comparison of the tetR codon usage to a mouse codon bias table. Values much lower than 1.0 are indicative of sequences which may translate poorly. For reference, a comparison of tetR to a Tobacco codon 20 bias table is included (light line). In transgenic tobacco, the tetR regulation system functions very efficiently, suggesting that for this gene, codon bias may be an important factor for efficient expression.

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7.1.7. COMPOSITIONAL ANALYSIS OF SYNTHETIC tetR
Figure 17 depicts the structure of the synthetic
tetR gene, now devoid of splice donor signal
sequences, with only a single splice acceptor signal
remaining (A). This is not the splice acceptor which
was active in the 345 construct. The percentage of G
and C bases has been significantly improved, while the
frequency of CpG base pairs has been kept to a
minimum. A CpG base pair is frequently the site for
DNA methylation, which can negatively effect the
expression of a gene. The codon bias of the synthetic

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tetR gene is vastly improved. The graph depicts the results when the synthetic tetR coding sequence is compared to the same mouse codon bias table used previously.

7.2 RESULTS

7.2.1. EXPRESSION OF tetR IN CONSTRUCT 345 OFFSPRING

construct was produced. The construct, referred to as Construct 345 is depicted in Figure 10. In the 345 construct the coding region of tetR is augmented with a nuclear localization signal sequence to increase the nuclear concentration of repressor. The tetR coding region was inserted into the first exon of the bGH gene. The bGH gene then acts as a genomic carrier, providing multiple introns, which may improve expression, and a strong polyadenylation signal, which may improve the processing and stability of the message.

The new repressor was coinjected with the bGH gene from construct 252. The resulting transgenic animals contain the new repressor, and a PEPCK regulated bGH gene with the tetR operators located just 3' of the PEPCK TATA-box element. Offspring of these animals were screened for bGH induction (FIG. 11). Of the lines tested only one, line 14, showed tetracycline dependent regulation of bGH, and in this one case there was still a significant base level of bGH expression. Northern analysis, performed to determine the levels of tetR mRNA expressed in the transgenic mice, indicated that the tetR gene was still not expressed at a high level.

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To detect tetR mRNA with higher sensitivity the tetR mRNA was analyzed using RNase protection. This technique revealed that the mRNA was shorter then

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expected (FIG. 12). Subsequent analysis using reverse transcriptase-PCR with primers that amplify the entire coding region of tetR confirmed that the mRNA was significantly shorter then expected (FIG. 13).

Sequence analysis of these RT-PCR products indicated that an unexpected splicing event had occurred. This splicing process occurred between a splice donor signal in the 5' end of the tetR coding region and a splice acceptor approximately 400 bp 3' of the start codon. The resulting mRNA is therefore deleted of the tetR DNA binding domain and about two third of the entire coding region. This mRNA could not possibly make a functional repressor.

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7.2.2. OPTIMIZATION OF tetR CONSTRUCT

A more detailed analysis of the tetR coding sequence indicated that the codons used in this gene are poorly suited for expression in mammalian cells (FIG. 14). Therefore, it appears that the inefficiency of the tetR system is the result of two processes: (i) aberrant splicing of the RNA to produce a nonfunctional message; and (ii) inefficient translation which can lead to rapid mRNA turnover.

To circumvent the problems of internal splicing and potential problems due to codon bias and G-C content, a synthetic tetR gene was designed. The components of the synthetic tetR gene were synthesized as four overlapping double stranded cassettes. Each cassette was cloned in puc19. The resulting plasmids designated pLT-1, pLT-2, pLT-3 and pLT-5, as depicted in Figure 15, have been deposited with ATCC and assigned accession numbers _____, ____, and _____, and _____, respectively. The synthetic tetR (syn-tetR) has been designed to encode exactly the same protein

product, but is devoid of splice signals and has

greatly improved codon usage for mammalian cells. The sequence of the of the syn-tetR is indicated in Figure 16. The predicted analysis for splicing signals, G+C content, and codon usage are depicted in Figure 17.

8. DEPOSIT OF MICROORGANISMS

The following microorganisms have been deposited with the American Type Culture Collection, (ATCC),

Rockville, Maryland and have been assigned the following accession numbers:

	<u>Microorganism</u>	Date of Deposit	Accession No.
	pLT-1	August 25, 1993	
•	pLT-2	August 25, 1993	•
15	pLT-3	August 25, 1993	
	pLT-5	August 25, 1993	
	pPCK_NbGH	August 25, 1993	

The present invention is not to be limited in scope by the microorganisms deposited since the deposited embodiments are intended as illustrations of single aspects of the invention and any microorganisms which are functionally equivalent are within the scope of the invention.

The present invention is not to be limited in scope by the exemplified embodiments which are intended as illustrations of single aspects of the invention, and any clones, DNA or amino acid sequences which are functionally equivalent are within the scope of the invention. Indeed, various modifications of the invention in addition to those described herein will become apparent to those skilled in the art from the foregoing description and accompanying drawings. Such modifications are intended to fall within the scope of the appended claims.

It is also to be understood that all base pair sizes given for nucleotides are approximate and are used for purposes of description.

Various publications are cited herein, which are hereby incorporated by reference in their entirety.

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- 40 -

SEQUENCE LISTING

(1) GENERAL INFORMATION:

- (i) APPLICANT: Byrne, Guerard
- (ii) TITLE OF INVENTION: TETRACYCLINE REPRESSOR-MEDIATED BINARY REGULATION SYSTEM FOR CONTROL OF GENE EXPRESSION IN TRANSGENIC ANIMALS
- (iii) NUMBER OF SEQUENCES: 15
- (iv) CORRESPONDENCE ADDRESS:
 - (A) ADDRESSEE: Pennie & Edmonds
 - (B) STREET: 1155 Avenue of the Americas
 - (C) CITY: New York
 - (D) STATE: New York
 - (E) COUNTRY: U.S.A.
 - (F) ZIP: 10036-2711
- (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.25
- (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER: US 07/935,763
 - (B) FILING DATE: 26-AUG-1992
 - (C) CLASSIFICATION:
- (viii) ATTORNEY/AGENT INFORMATION:
 - (A) NAME: Coruzzi, Laura A.
 - (B) REGISTRATION NUMBER: 30,742
 - (C) REFERENCE/DOCKET NUMBER: 6794-025
 - (ix) TELECOMMUNICATION INFORMATION:
 - (A) TELEPHONE: 212 790-9090
 - (B) TELEFAX: 212 869-8864/9741 (C) TELEX: 66141 PENNIE
- (2) INFORMATION FOR SEQ ID NO:1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 59 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: unknown
 - (ii) MOLECULE TYPE: DNA (genomic)
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

TTGACACTCT ATCATTGATA GAGTTATTTT ACCACTCCCT ATCAGTGATA GAGAAAAGT

59

(2) INFORMATION FOR SEQ ID NO:2:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 70 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: unknown

(11) NODDOUD IIIB. DIA (GENOMIC)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:	
GAATTCGATA CTCTATCATT GATAGAGTAT CAAGCTTATC CCTATCAGTG ATAGAGATAC	60
CGTCGACCTC	70
(2) INFORMATION FOR SEQ ID NO:3:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 49 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: unknown	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:	
ACTCTATCAT TGATAGAGTT ACTATTTAAA TCCCTATCAG TGATAGAGA	49
(2) INFORMATION FOR SEQ ID NO:4:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 71 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: unknown	
(ii) MOLECULE TYPE: DNA (genomic)	-
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4: GGAATTCGAT ACTCTATCAT TGATAGAGTA TCAAGCTTAT CCCTATCAGT GATAGAGATA	60
	60
CCGTCGACCT C	71
(2) INFORMATION FOR SEQ ID NO:5:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 624 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: unknown 	
(ii) MOLECULE TYPE: DNA (genomic)	
(ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 1624	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:	
ATG TCT AGA TTA GAT AAA AGT AAA GTG ATT AAC AGC GCA TTA GAG CTG Met Ser Arg Leu Asp Lys Ser Lys Val Ile Asn S.r Ala Leu Glu Leu 1 5 10 15	48
CTT AAT GAG GTC GGA ATC GAA GGT TTA ACA ACC CGT AAA CTC GCC CAG	96

Leu	Asn	Glu	Val 20	Gly	Ile	Glu	Gly	Leu 25	Thr	Thr	Arg	Lys	Leu 30	Ala	Gln	
AAG Lys	CTA Leu	GGT Gly 35	GTA Val	GAG Glu	CAG Gln	CCT Pro	ACA Thr 40	TTG Leu	TAT Tyr	TGG Trp	CAT His	GTA Val 45	Lys	AAT Asn	AAG Lys	14
CGG Arg	GCT Ala 50	TTG Leu	CTC Leu	GAC Asp	GCC Ala	TTA Leu 55	GCC Ala	ATT Ile	GAG Glu	ATG Met	TTA Leu 60	GAT Asp	AGG Arg	CAC His	CAT His	19
ACT Thr 65	CAC His	TTT Phe	TGC Cys	CCT Pro	TTA Leu 70	GAA Glu	GGG Gly	GAA Glu	AGC Ser	TGG Trp 75	CAA Gln	GAT Asp	TTT Phe	TTA Leu	CGT Arg 80	24
AAT Asn	AAC Asn	GCT Ala	AAA Lys	AGT Ser 85	TTT Phe	AGA Arg	TGT Cys	GCT Ala	TTA Leu 90	CTA Leu	AGT Ser	CAT His	CGC Arg	GAT Asp 95	GGA Gly	28
GCA Ala	AAA Lys	GTA Val	CAT His 100	TTA Leu	GGT Gly	ACA Thr	CGG Arg	CCT Pro 105	ACA Thr	GAA Glu	AAA Lys	CAG Gln	TAT Tyr 110	GAA Glu	ACT Thr	330
CTC Leu	GAA Glu	AAT Asn 115	CAA Gln	TTA Leu	GCC Ala	TTT Phe	TTA Leu 120	TGC Cys	CAA Gln	CAA Gln	GGT Gly	TTT Phe 125	TCA Ser	CTA Leu	GAG Glu	384
AAT Asn	GCA Ala 130	TTA Leu	TAT Tyr	GCA Ala	CTC Leu	AGC Ser 135	GCT Ala	GTG Val	GGG Gly	CAT His	TTT Phe 140	ACT Thr	TTA Leu	GGT Gly	TGC Cys	432
GTA Val 145	TTG Leu	GAA Glu	GAT A sp	CAA Gln	GAG Glu 150	CAT His	CAA Gln	GTC Val	GCT Ala	AAA Lys 155	GAA Glu	GAA Glu	AGG Arg	GAA Glu	ACA Thr 160	480
CCT Pro	ACT Thr	ACT Thr	GAT Asp	AGT Ser 165	ATG Met	CCG Pro	CCA Pro	TTA Leu	TTA Leu 170	CGA Arg	CAA Gln	GCT Ala	ATC Ile	GAA Glu 175	TTA Leu	528
TTT Phe	GAT Asp	CAC His	CAA Gln 180	GGT Gly	GCA Ala	GAG Glu	CCA Pro	GCC Ala 185	TTC Phe	TTA Leu	TTC Phe	GGC Gly	CTT Leu 190	GAA Glu	TTG Leu	576
ATC Ile	ATA Ile	TGC Cys 195	GGA Gly	TTA Leu	GAA Glu	AAA Lys	CAA Gln 200	CTT Leu	AAA Lys	TGT Cys	GAA Glu	AGT Ser 205	GGG Gly	TCT Ser	TAA	624

(2) INFORMATION FOR SEQ ID NO:6:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 207 amino acids(B) TYPE: amino acid

 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:

Met Ser Arg Leu Asp Lys Ser Lys Val Ile Asn Ser Ala Leu Glu Leu

Leu Asn Glu Val Gly Ile Glu Gly Leu Thr Thr A-c Lys Leu Ala Gln

Lys Leu Gly Val Glu Gln Pro Thr Leu Tyr Trp His Val Lys Asn Lys

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Arg	Ala 50	Leu	Leu	Asp	Ala	Leu 55	Ala	Ile	Glu	Met	Leu 60	Asp	Arg	His	His		
Thr 65	His	Phe	Сув	Pro	Leu 70	Glu	Gly	Glu	Ser	Trp 75	Gln	Asp	Phe	Leu	Arg 80		
Asn	Asn	Ala	Lys	Ser 85	Phe	Arg	Сув	Ala	Leu 90	Leu	Ser	His	Arg	Asp 95	Gly		
Ala	Lys	Val	His 100	Leu	Gly	Thr	Arg	Pro 105	Thr	Glu	Lys	Gln	Туг 110	Glu	Thr		
Leu	Glu	Aøn 115	Gln	Leu	Ala	Phe	Leu 120	Сув	Gln	Gln	Gly	Phe 125	Ser	Leu	Glu		
Asn	Ala 130	Leu	Tyr	Ala	Leu	Ser 135	Ala	Val	Gly	His	Phe 140	Thr	Leu	Gly	Сув		
Val 145	Léu	Glu	Авр	Gln	Glu 150	His	Gln	Val	Ala	Lys 155	Glu	Glu	Arg	Glu	Thr 160		
Pro	Thr	Thr	Asp	Ser 165	Met	Pro	Pro	Leu	Leu 170	Arg	Gln	Ala	Ile	Glu 175	Leu		
Phe	Asp	His	Gln 180	Gly	Ala	Glu	Pro	Ala 185	Phe	Leu	Phe	Gly	Leu 190	Glu	Leu		
Ile	Ile	Cys 195	Gly	Leu	Glu	Lys	Gln 200	Leu	Lys	Сув	Glu	Ser 205	Gly	Ser			
(2)	(i)) SE(() () ()	QUENCA) LI B) TI C) SI	CE CI ENGTI YPE: TRANI DPOLO	SEQ HARACH: 92 nucl DEDNI DGY: (PE:	CTER: 2 bas leic ESS: unks	ISTIC se pa acio sino nown	CS: airs i gle	c)								
	(xi) SEG	QUENC	CE DI	ESCR:	IPTIC	ON: S	SEQ :	ID N	0:7:							
CGG	CCCT	ATA Z	AAAA	GCGAI	AG CO	GCGC	GCG	G GC	GGGA	GTCG	CTG	CGTT	GCC '	TTCG	ccccg1	r	60
GCC	CCGC	rcc (GCGC	CGCC:	IC G	CGCC	GCCC	G CC									92
(2)	INF	ORMA'	TION	FOR	SEQ	ID I	8 : ON	:									
	(i	() ()	A) L1 B) T1 C) S1	ENGTI YPE : IRANI	HARAG H: 6: nuc: DEDNI DGY:	l bas leic ESS:	se pa acio sin	airs d									
	(ii) MO	LECU	LE T	YPE:	DNA	(ge	nomi	c)								
	(xi) SE	QUEN	CE D	ESCR	IPTI(ON:	SEQ	ID N	0:8:							
AAG.	AAGT:	ATA '	TTAG	AGCG	AG T	CTTT	CTGC.	A CA	CACG	ATCA	ССТ	TTCC	TAT	CAAC	CCCAC	r	60

(2) INFORMATION FOR SEQ ID NO:9:

	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 74 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: unknown	
	(ii) MOLECULE TYPE: DNA (genomic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:	
GTAT	TATGTT TTATGTTACT GTAAAAGATG TAAAGAGAGG CACGTGGTTA AGCTCTCGGG	60
GTGT	GGACTC CACC	74
(2)	INFORMATION FOR SEQ ID NO:10:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 73 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: unknown	
	(ii) MOLECULE TYPE: DNA (genomic)	
•	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:	
cccc	CCAAGC ATAAACCCTG GCGCGCTCGC GGCCCGGCAC TCTTCTGGTC CCCACAGACT	60
CAGA	GAGAAC CCA	73
(2)	INFORMATION FOR SEQ ID NO:11:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 74 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: unknown	
	(ii) MOLECULE TYPE: DNA (genomic)	
	(*i) SEQUENCE DESCRIPTION: SEQ ID NO:11:	-
TAGG	CAGCAG GCATATGGGA TGGGATATAA AGGGGCTGGA GCACTGAGAG CTGTCAGAGA	60
TTTC	TCCAAC CCAG	74
(2)	INFORMATION FOR SEQ ID NO:12:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 19 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: unknown	
	(ii) MOLECULE TYPE: DNA (genomic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:	

ACTCTATCAT TGATAGAGT	19
2) INFORMATION FOR SEQ ID NO:13:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 19 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: unknown	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:	
ACTCTATCAA TGATAGAGT	19
(2) INFORMATION FOR SEQ ID NO:14:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 19 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: unknown	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:	
TCCCTATCAG TGATAGAGA	19
(2) INFORMATION FOR SEQ ID NO:15:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 19 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: unknown	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:15:	
TCTCTATCAC TGATAGGGA	19

-46International Application No: PCT/

MICROC	PRGANISMS
Optional Sheet in connection with the microorganism re	ferred to on page <u>38</u> , lines <u>7-23</u> of the description '
A. IDENTIFICATION OF DEPOSIT	
Further deposits are identified on an additional sheet	•
Name of depositary institution '	
American Type Culture Collection	
j C	
Address of depositery institution (including postal	code and country) *
12301 Perklawn Drive Rockville, MD 10582 US	
Date of deposit ' August 25, 1993 Accession No.	umber * <u>N/A</u>
B. ADDITIONAL INDICATIONS · (leave black if not applicable	ole). This information is continued on a separate attached sheet 13
C. DESIGNATED STATES FOR WHICH INDICATIO	UNS ARE MADE " (If the indication we are the indication)
D. SEPARATE FURNISHING OF INDICATIONS ' (les	an black if no analisable
The indications listed below will be submitted to the International B	
"Accession Number of Deposit")	умово нест пороску на умама насел от на импевона с.д.,
E. This sheet was received with the International app	lication when filed (to be checked by the receiving Office)
- 3. · · · · · · · · · · · · · · · · · ·	(Authorized Officer) Lelrly
☐ The date of receipt (from the applicant) by the Int	ternational Bureau *
was	(Authorized Officer)
	(FIGURE OFFICE)

Form PCT/RO/134 (January 1981)

International Application No: PCT/

Form PCT/RO/134 (cont.)

American Type Culture Collection

12301 Parkiawn Drive Rockville, MD 10582 US

Accession No.	Date of Deposit
N/A	August 25, 1993

- nucleic acid mol cul comprising an animal promoter
 selement that comprises a tetR operator sequence.
 - 2. The nucleic acid molecule of claim 1 in which the tetR operator sequence is positioned 3' to a TATA-box sequence.

- 3. The nucleic acid molecule of claim 1 in which the promoter element is the PEPCK promoter.
- 4. The nucleic acid molecule of claim 3 in which the tetR operator sequence has been inserted into the NheI site of the PEPCK promoter element.
- 5. The nucleic acid molecule of claim 1, 2, 3 or 4 in which the promoter element controls the expression of a gene of interest.
 - 6. The nucleic acid molecule of claim 5 in which the gene of interest is bovine growth hormone.
- 7. A non-human transgenic animal that carries, as a transgene, the nucleic acid molecule of claim 1, 2, 3 or 4.
- A non-human transgenic animal that carries,
 as a transgene, the nucleic acid molecule of claim 5.
 - 9. A non-human transgenic animal that carries, as a transgene, the nucleic acid molecule of claim 6.

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10. The non-human transgenic animal of claim 7 that further carries a transgene encoding the tetR repressor protein.

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- 11. The non-human transgenic animal of claim 8 that further carries a transgene encoding the tetR repressor protein.
- 12. The non-human transgenic animal of claim 9 that further carries a transgene encoding the tetR repressor protein.
- 13. A non-human transgenic animal that carries a transgene encoding the tetR repressor protein.
 - 14. A method of selectively inducing the expression of a gene of interest in a non-human transgenic animal comprising administering a tetracycline compound to a non-human transgenic animal that carries a first transgene which is a gene of interest under the control of a promoter element modified to comprise a tetR operator sequence and a second transgene encoding the tetR repressor protein.

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- 15. A non-human transgenic animal that carries (i) a first transgene that encodes bovine growth hormone and is under the control of PEPCK promoter element modified to contain a tetR operator at the NheI site; and (ii) a second transgene that encodes tetR repressor protein.
- 16. The transgenic animal of claim 15 that is a mouse.

nucleic acid molecule comprising an optimized tetR gene as depicted in Figure 16.

- 19. The non-human transgenic animal of claim 7

 10 that further carries an optimized transgene encoding the tetR repressor protein and having a sequence as depicted in Figure 16.
- 20. The non-human transgenic animal of claim 8

 15 that further carries an optimized transgene encoding
 the tetR repressor protein and having a sequence as
 depicted in Figure 16.
- 21. The non-human transgenic animal of claim 9
 20 that further carries an optimized transgene encoding
 the tetR repressor protein and having a sequence as
 depicted in Figure 16.
- 22. A non-human transgenic animal that carries
 25 an optimized transgene encoding the tetR repressor
 protein and having a sequence as depicted in Figure
 16.
- 23. A method of selectively inducing the

 25 expression of a gene of interest in a non-human

 26 transgenic animal comprising administering a

 27 tetracycline compound to a non-human transgenic animal

 28 that carries a first transgene which is a gene of

 29 interest under the control of a promoter element

 29 modified to comprise a tetR operator sequence and a

 20 second optimized transgene encoding the tetR repressor

protein and having a sequence as depicted in Figure 16.

- 24. A non-human transgenic animal that carries

 (i) a first transgene that encodes bovine growth
 hormone and is under the control of PEPCK promoter
 element modified to contain a tetR operator at the
 NheI site; and (ii) a second optimized transgene that
 encodes tetR repressor protein that has a sequence as
 depicted in Figure 16.
 - 25. The transgenic animal of claim 24 that is a mouse.
- 26. The transgenic animal of claim 24 that is a pig.

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FIEURE

Tn10

Operator 1

Operator sites in Tn10.

Operator 2

TTGACACTCTATCATTGATAGAGTTATTTTACCACTCCCTATCAGTGATAGAGAAAGT

Oligonucleotides

Hind3

Operator 1

EcoR1

Operator 2

General purpose operator.

GAATTCGAT ACTCTATCATTGATAGAGT ATCAAGCTTA TCCCTATCAGTGATAGAGA TACCGTCGACCTC

PEPCK-TATA box operator.

Operator 1

Operator 2.

ACTCTATCATTGATAGAGT TACTATTTAAA TCCCTATCAGTGATAGAGA

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		* * *					
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EcoR1 OP1 linker
ggaattcgat-ACT CTA TCA TTG ATA GAG TAT CAA GCT TAT CCC

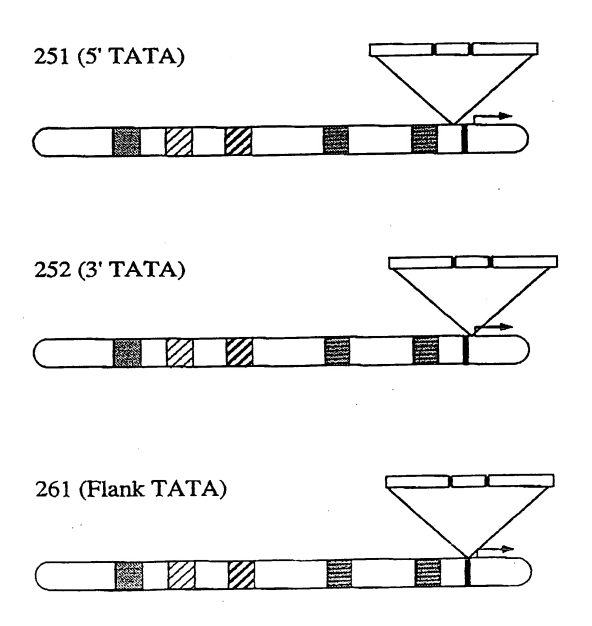
OP2 AccI
TAT CAG TGA TAG AGA-taccgtcgacctc

FIGURE 18.

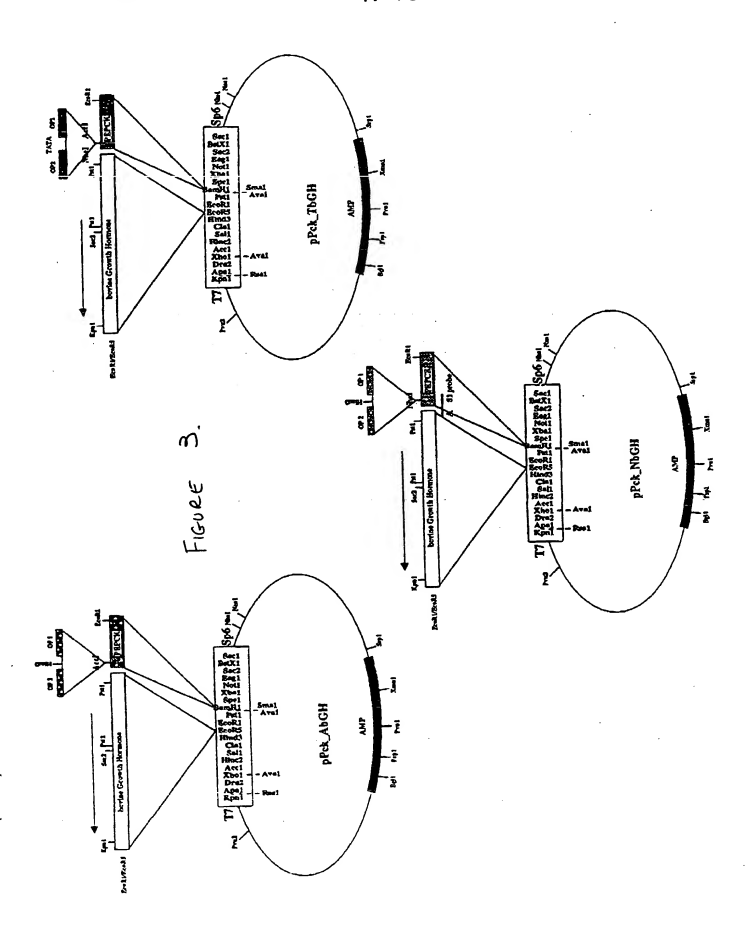
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FIGURE 2.

MODIFIED PEPCK PROMOTERS



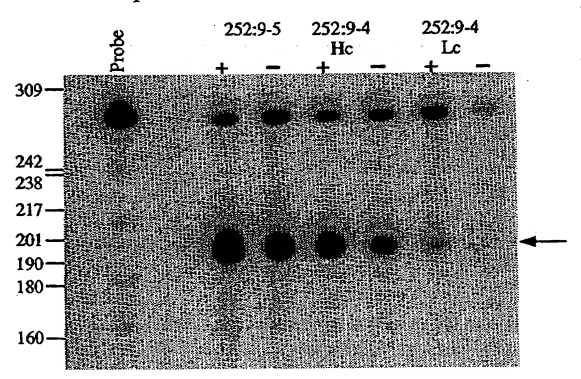
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FIGURE 4.

S₁ Nuclease Protection: 5' Start Site



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2.1124								7.			
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				•							
			•								

FIGURE 5.

10		20	30 *		40
ATG TCT AGA TO TETRACYCLINGbbbca_1520a	L D K NE REPRESS	S K OR PROTE:	DN (TETR)	CODON_ST	ART=1> _bb>
50	60 *	•	70	80 *	90 *
CTG CTT AAT G L L NTETRACYCLIbbb <1480_aa14	AG GTC GGA E V G NE REPRESS	OR PROTE	IN (TETR) RNA (SPLI	; CODON_ST	ART=1> _bb>
100		110	120		130
GCC CAG AAG C A Q KTETRACYCLIb_b_b <a_1430_a< td=""><td>L G V NE REPRESS</td><td>E Q SOR PROTE</td><td>IN (TETR)</td><td>CODON ST</td><td>ART=1> _bb></td></a_1430_a<>	L G V NE REPRESS	E Q SOR PROTE	IN (TETR)	CODON ST	ART=1> _bb>
140	150		.60 *	170	180
AAA AAT AAG (K N K	R A L INE REPRES	SOR PROTE	EIN (TETR); CODON_S	TART=1> _bb>
190		200	210		220
	H T H	SOR PROT	ein (Tetr); CODON_S	TART=1>
230	240		250 *	260 *	270 *
	INE REPRES	SOR PROT	EIN (TETF	R); CODON_S	TART=1>
28		290	306		310
TETRACYCI	R D C	g a i SSOR PRO Eddersor	C V H PEIN (TET) MRNA (SP)	R); CODON_S	ACA CGG CCT T R P> START=1>b_b> 0aa

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	market in

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FIGURE 5 (continued)

320	3	30	340		350 *	. 30	60 *
T E	CYCLINE R	WD BEBEE E <u>beeseop</u> Y E T	L E PROTEIN (SOR MRNA	n ((TETR) ; (SPLIT	CODON	TART=1 _ bb_	
		200		390		400	
	370 *	380		*	m.m C	*	·~
C Q TETRA	CAA GGT TI Q G F CYCLINE RE 	EPRESSOR F	PROTEIN (TETR);	CODON_S _bb_ _a_1130	TART=1b_b_ _a_a_a	> >
410		20	430 *		440 *	45	•
A V	GGG CAT TO G H I CYCLINE RO 	epressor 1	PROTEIN (TETR);	CODON	TART=1	
	460	470		480		490 *	
H Q TETRA	GTC GCT A V A ACYCLINE R D	K E E EPRESSOR :	PROTEIN ((TETR);	CODON	START=1bb_	>
500	5	10	520		530 *	5	40
M P	CCA TTA T P L ACYCLINE F D D TF D 1020_5	L R Q EPRESSOR	PROTEIN	(TETR);	CODON	START=1_ b_ b_	<u>`</u>
	550	560	_	570 *		580 *	
G A TETR	GAG CCA (E P ACYCLINE b_b_T 80_a_a	A F	PROTEIN	(TETR):	CODONL	START=1_b	> >
590		600	610		620		
G L TETF	GAA AAA E K VACYCLINE bTETR a903	Q L REPRESSOR	AA TGT GI K C I PROTEIN	(TETR)	GGG TCT G S ; CODON	> b>	

		 9 ;
·		
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		!

FIGURE

(Non-specific expression in most tissues)

Kost TA, Theodorakis N, and Hughes SH (1983) Nucleic Acids Res. 11; 8287 - 8301.

AAGAAG<u>TATA</u>GAGCGAGTCTTTCTGCACACGCATCACCTTTCCTATCAACCCCACTA Albumin

Gorski K, Carneiro M, and Schibler U. (1986) Cell 62; 991 - 998. specific)

GTATTAT<u>GTITTATG</u>TTACTGTAAAAGATGTAAAGAGAGCACGTGGTTAAGCTCTOGGGGTGTGGACTCCACC Human CD-2

Lang G, Wotton D, Owen MJ, Sewell WA, Brown MH, Mason DY, Crumpton MJ, and Kioussis D. (1988) EMBO J. 7; 1675

COCCCCAAGC<u>AİAAA</u>ČCCTGGCGCGCGCCCGGCCCGCCACTCTTCTGGTCCCCACAGACTCAGAGAGAACCCA alpha-globin *

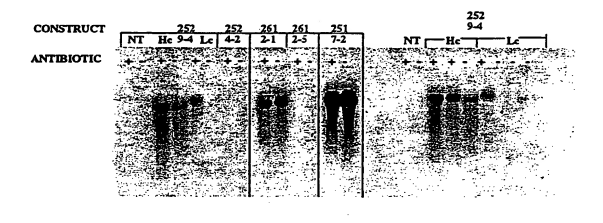
Liebhaber SA, Goossens MJ, and Wai Kan Y. (1980) Proc. Natl. Acad. Sci. USA. 77; 7054 - 7058. (Red blood cells)

TAGGCAGCAGCATATÓGGATGGG<u>ATÁTAAA</u>GGGGCTGGAGCACTGAGAGCTGTCAGAGATTTCTCCAACCCAG Mouse Cardiac Myosin Heavy Chain

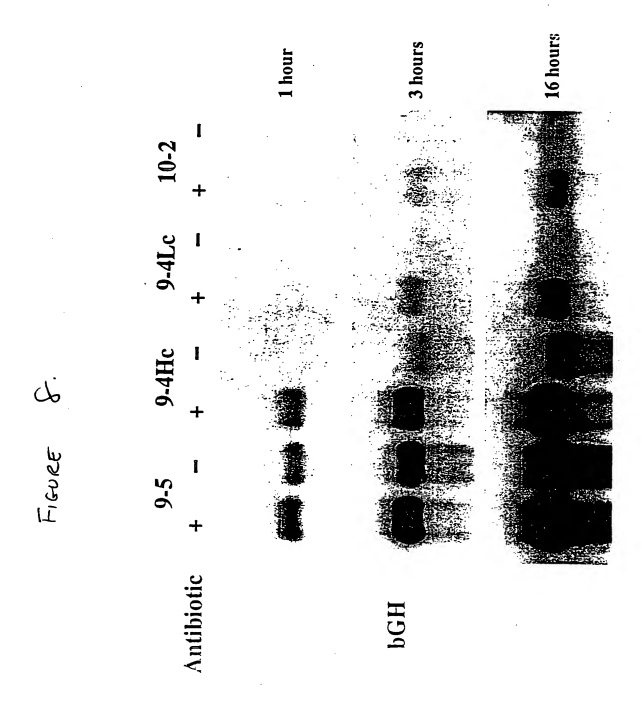
- 998. Tanigawa G, Jarcho JA, Kass S, Solomon SD, Vosberg H.-P, Seidman JG, and Seidman CE. (1990) Cell 62; 991

FIGURE 7.

INDUCTION OF BOVINE GROWTH HORMONE mRNA BY TETRACYCLINE

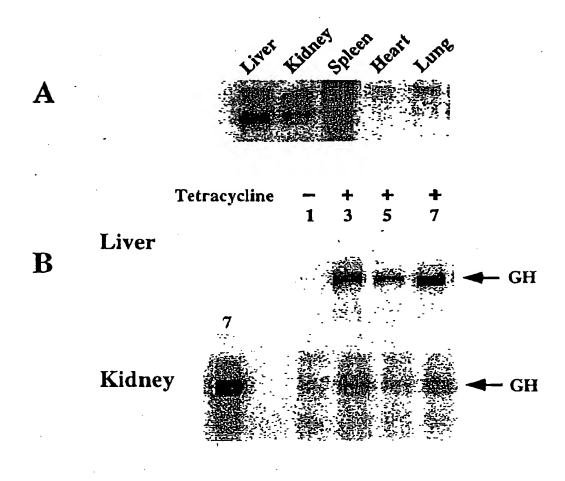


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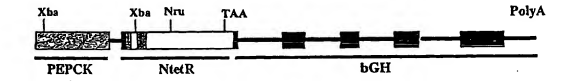
Tissue Specificity. and Tetracycline Induction of bGH in Line 10-2



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Figure 10

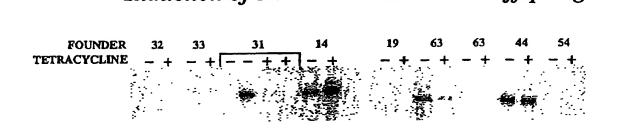
345 Repressor Construct



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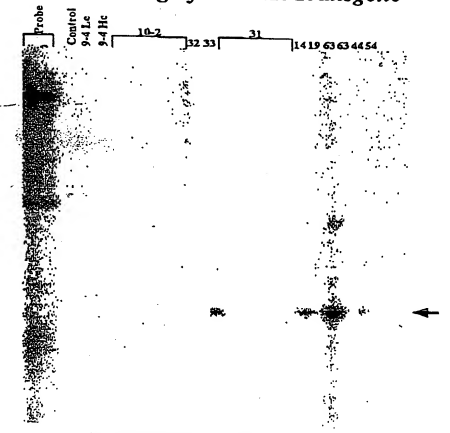
Figure 11

Induction of bGH in Construct 345 Offspring

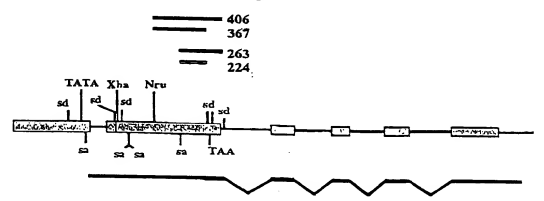


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Expression and Alternative Processing of the tetR Transgene

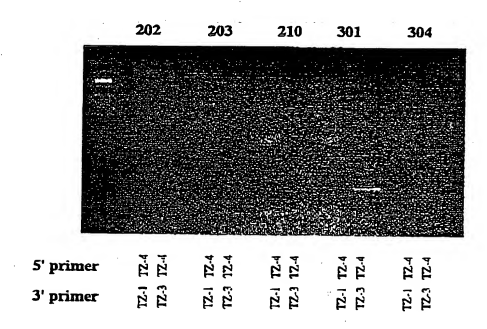


Protected fragments



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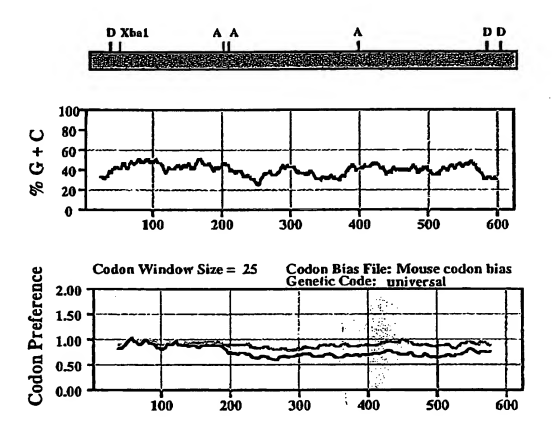
5' Structure of tetR mRNA



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Composition Analysis of Wild Type Tn10 tetR Gene.



Synthetic tetR Component Sequences

LT-1

EcoR5 and EcoR1 GATATCGAATTCATGAGTAGATTGGACAAGAGCAAAGTGATCAATAGTGC TCTGGAGCTGTTGAATGAAGTGGGCATAGAAGGTCTGACTACCAGAAAGC

AATAAGAGGGCTCTGCTGGATGCATTGGCGGTACCAGGC - رايو لومه کام د Nsi1 Kpnl

LT-2

Kpnl Nsi1

ACACTTCTGCCCACTGGAAGGCGAGAGTTGGCAGGACTTCCTGAGGAACA ATGCTAAGAGTTTCAGATGTGCTCTGTTGAGCCACAGAGACGGTGCTAAA **GTGCAC**CTGGAATTCGAGC ApaL1

TGGCCCAGAAGCTGGGAGTGGAGCAGCCAACATTGTACTGGCATGTGAAG

EcoR1

LT-3

EcoR1 ApaL1 GCTCGAATTCAAAGTGCACCTGGGTACAAGGCCAACAGAGAAACAGTACG AGACCCTGGAGAACCAGCTGGCATTTCTGTGCCAACAAGGCTTCAGCCTG GAGAATGCATTGTATGCTCTGAGTGCTGTGGGTCACTTCACACTGGGTTG TCTCCTGGAGGACCAGGAGCACCAGGTGGCCAAGGAGGAGAGGGAGACCC CAACCACTGACAGCATGCCCCGGATCCGAGC Sph1 BanH1

LT-5

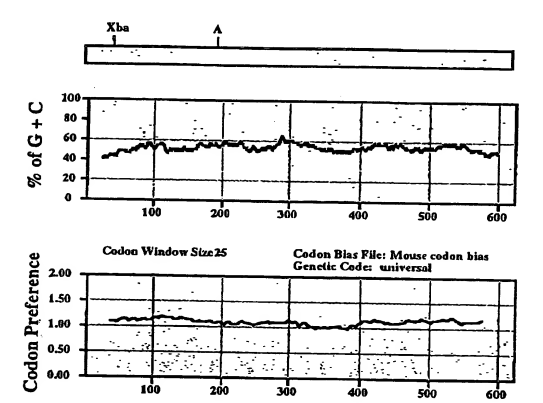
BamH1 Sph1 GCTCGGATCCACAGCATGCCCCCATTGCTGAGACAGGCCTATGAGCTGTT TGACCACCAAGGGGCAGAGCCTGCTTTTCTGTTTGGCCTGGAGCTCATCA TCTGTGGTCTGGAGAGCAGCTGAAGTGTGAGAGTGGCTCCTGAAGCTTG Hind3/EcoR5 ATATC

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Sequence of Synthetic tetR Gene.

TCATGAGTAG	ATTGGACAAG	AGCAAAGTGA
TCTGGAGCTG	TTGAATGAAG	TGGGCATAGA
ACCAGAAAGC	TGGCCCAGAA	GCTGGGAGTG
CATTGTACTG	GCATGTGAAG	AATAAGAGGG
TGCATTGGCC	ATTGAGATGC	TGGACAGACA
TTCTGCCCAC	TGGAAGGCGA	GAGTTGGCAG
GGAACAATGC	TAAGAGTTTC	AGATGTGCTC
CAGAGACGGT	GCTAAAGTGC	ACCTGGGTAC
GAGAAACAGT	ACGAGACCCT	GGAGAACCAG
TGTGCCAACA	AGGCTTCAGC	CTGGAGAATG
TCTGAGTGCT	GTGGGTCACT	TCACACTGGG
GAGGACCAGG	AGCACCAGGT	GGCCAAGGAG
CCCCAACCAC	TGACAGCATG	CCCCCATTGC
CATAGAGCTG	TTTGACCACC	AAGGGGCAGA
CTGTTTGGCC	TGGAGCTCAT	CATCTGTGGT
AGCTGAAGTG	TGAGAGTGGC	TCCTGAAGCT
	•	
	TCTGGAGCTG ACCAGAAAGC CATTGTACTG TGCATTGGCC TTCTGCCCAC GGAACAATGC CAGAGACGGT GAGAAACAGT TGTGCCAACA TCTGAGTGCT GAGGACCAGG CCCCAACCAC CATAGAGCTG CTGTTTGGCC	TCTGGAGCTG TTGAATGAAG ACCAGAAAGC TGGCCCAGAA CATTGTACTG GCATGTGAAG TGCATTGGCC ATTGAGATGC TTCTGCCCAC TGGAAGGCGA GGAACAATGC TAAGAGTTTC CAGAGACGGT GCTAAAGTGC GAGAAACAGT ACGAGACCCT TGTGCCAACA AGGCTTCAGC TCTGAGTGCT GTGGGTCACT GAGGACCAGG AGCACCAGGT CCCCAACCAC TGACAGCATG CATAGAGCTG TTTGACCACC CTGTTTGGCC TGGAGCTCAT

Compositional analysis of Synthetic tetR



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INTERNATIONAL SEARCH REPORT

International application No. PCT/US93/08230

	SSIFICATION OF SUBJECT MATTER					
	:C12N 15/00; C07H 21/00 :800/2; 435/172.3, 320.1; 536/27					
	to International Patent Classification (IPC) or to both	national classificati n and IPC	-			
B. FIEL	LDS SEARCHED					
Minimum d	ocumentation searched (classification system followe	d by classification symbols)				
U.S. :	800/2; 435/172.3, 320.1; 536/27; 514/152; 935/40,	43, 111				
Documentat	tion searched other than minimum documentation to the	e extent that such documents are included	in the fields searched			
		•				
Electronic d	lata base consulted during the international search (na	ame of data base and, where practicable,	, search terms used)			
Please Sec	e Extra Shect.					
C. DOC	UMENTS CONSIDERED TO BE RELEVANT					
Category*	Citation of document, with indication, where a	ppropriate, of the relevant passages	Relevant to claim No.			
X	US,A, 5,075,229 (Hanson et al.) iss	ued 24 December 1991, see	<u>7-9</u> 1-6,			
Y	entire document.		10-26			
Y,P	US,A, 5,221,778 (Byrne et al.) issued 22 June 1993, see entire 1-26 document.					
Y	Mol. Gen. Genet., Volume 227, Numlet al., Regulating a modified CaMV encoded Tet repressor in transgenic tobentire document.	35S promoter by the Tn10-	1-26			
	·					
X Furth	er documents are listed in the continuation of Box C	. See patent family annex.				
"A" doc	ocial categories of cited documents: nument defining the general state of the art which is not considered	"T" Inter document published after the inter date and not in conflict with the applic principle or theory underlying the inv	stion but cited to understand the			
	to part of particular relovance tier document published on or after the international filing date	"X" document of particular relevance; the	o claimed invention cannot be			
"L" doc	nument which may throw doubts on priority claim(s) or which is d to establish the publication date of another citation or other	considered novel or cannot be conside when the document is taken alone	•			
spo	cial remen (as specified) nument referring to an oral disclosure, use, exhibition or other	"Y" document of particular relevance; the considered to involve an inventive combined with one or more other such being obvious to a person skilled in the person skilled in the person skilled in the person skilled in the person skilled in the person skilled in person step when the document is a documents, such combination				
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	nailing address of the ISA/US	Authorized flicer	Mr. Don			
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INTERNATIONAL SEARCH REPORT

International application No. PCT/US93/08230

C (Continue	ntion). DOCUMENTS CONSIDERED TO BE RELEVANT	
Category*	Citation of document, with indication, where appropriate, of the relevant passage	s Relevant to claim No
Y	The Plant Journal, Volume 2, Number 3, issued 1992, Gatz et "Stringent repression and homogeneous de-repression by tetracycline of a modified CaMV 35S promoter in intact transgetobacco plants", pages 397-404, see the entire document.	.,
Y	Proc. Natl. Acad. Sci. USA, Volume 89, issued June 1992, Gossen et al., "Tight control of gene expression in mammalian cells by tetracycline-responsive promoters", pages 5547-5551, sthe entire document.	1-26 ee
Y	Proc. Natl. Acad. Sci. USA, Volume 86, issued July 1989, Byret al., "Multiplex gene regulation: a two-tiered approach to transgene regulation in transgenic mice", pages 5473-5477, see entire document.	
Y	Proc. Natl. Acad. Sci. USA, Volume 88, issued February 1991 Ornitz et al., "Binary system for regulating transgene expression in mice: targeting int-2 gene expression with yeast <u>GALA/UAS</u> control elements", pages 698-702, see the entire document.	n
Y	Proc. Natl. Acad. Sci. USA, Volume 85, issued March 1988, Gatz et al., "Tn10-encoded tet repressor can regulate an operate containing plant promoter", pages 1394-1397, see the entire document.	1-26
Y	Nucleic Acids Research, Volume 16, Number 4, issued 1988, Khillan et al., "Gene transactivation mediated by the TAT gene human immunodeficiency virus in transgenic mice", pages 1423 1430, see the entire document.	
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INTERNATIONAL SEARCH REPORT

International application No. PCT/US93/08230

	ervations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This internat	ional report has not been established in respect f certain claims under Article 17(2)(a) for the following reasons:
	aims Nos.: cause they relate to subject matter not required to be searched by this Authority, namely:
ليا ل	laims Nos.: cause they relate to parts of the international application that do not comply with the prescribed requirements to such extent that no meaningful international search can be carried out, specifically:
	laims Nos.: cause they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
lox II Obs	servations where unity of invention is lacking (Continuation of item 2 of first sheet)
This Interna	tional Scarching Authority found multiple inventions in this international application, as follows: (Telephone Practice) Group I, claims 1-6 and 18, drawn to a nucleic acid molecule, classified in Class 536, subclass 27, for example.
subcl	Group II, claims 7-17 and 19-26, drawn to transgenic animals and a method of using the same, classified in Class 80 ass 2, for example.
ol A 🔲 ـا	s all required additional search fees were timely paid by the applicant, this international search report covers all searchable aims. s all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment any additional fee.
	s only some of the required additional search fees were timely paid by the applicant, this international search report cover the search report coverable that the search fees were paid, specifically claims Nos.:
	o required additional search fees were timely paid by the applicant. Consequently, this international search report stricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark on	Protest The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.

international application

PCT/US93/08230

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Electronic data bases consulted (Name of data base and where practicable terms used):

DIALOG (files 154, 55, 312)

U.S. Automated Patent System (file USPAT, 1975-1993)

Search terms: tetracycline, tetR, operator, repressor, binary, transactivate, PEPCK, 6GH, transgenic, mice, transresponder, Tn10, tet, inventor's name.

Form PCT/ISA/210 (extra sheet)(July 1992)*